CLAIMS



- 1. In a method which calls for administration of IFN-α, the improvement comprising coadministering an effective amount of an isolated immunostimulatory nucleic acid.
- 2. The improvement of claim 1, wherein the IFN- α is administered at a dose below the clinically established effective dose for IFN- α alone.
- 3. The improvement of claim 1, wherein the IFN-α is administered at the maximum tolerated dose for IFN-α in the absence of the nucleic acid.
 - 4. The improvement of claim 1, wherein the IFN- α is administered at least 20 percent below the maximum tolerated dose of IFN- α in the subject.
- 5. The improvement of claim 1, wherein the IFN-α is administered at least 30 percent below the maximum tolerated dose of IFN-α in the subject.
 - 6. The improvement of claim 1, wherein the IFN- α is administered at least 40 percent below the maximum tolerated dose of IFN- α in the subject.

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- 7. The improvement of claim 1, wherein the IFN- α is administered at least 50 percent below the maximum tolerated dose of IFN- α in the subject.
- 8. The improvement of claim 1, wherein the immunostimulatory nucleic acid is modified.
 - 9. The improvement of claim 1, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.

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- 10. The improvement of claim 1, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.
- 11. The improvement of claim 1, wherein the immunostimulatory nucleic acid is not a palindrome.
 - 12. The improvement of claim 1, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
- 13. The improvement of claim 1, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.
 - 14. The improvement of claim 13, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
 - 15. The improvement of claim 13, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.
 - 16. The improvement of claim 1, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, T-rich nucleic acids, and poly-G nucleic acids.
 - 17. The improvement of claim 1, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.
 - 18. The improvement of claim 1, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.
 - 19. The improvement of claim 1, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

ggGGTCAACGTTGAgggggGGtcgtcgttttgtcgttttgtcgtttgggggg

ODN 1585 SEQ ID NO:1 ODN 2022 SEQ ID NO:2 ODN 2184 SEQ ID NO:3

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tcgtcgttttgtcgttttgggggg	ODN 2185	SEQ ID NO:
ggggtcgacgtcgaggggg	ODN 2192	SEQ ID NØ:5
ggggtcatcgatgaggggg	ODN 2204	SEQ ID NO:6
ggGGGACGATCGTCgggggG	ODN 2216	SEQ LO NO:7
gggggtcgtacgacgggggg	ODN 2217	SEQ ID NO:8
ggGGACGATATCGTCgggggG	ODN 2245	SEQ ID NO:9
ggGGACGACGTCGTCgggggG	ODN 2246	SEQ ID NO:10
ggGGACGAGCTCGTCgggggG	ODN 2247 /	SEQ ID NO:11
ggGGGACGTACGTCgggggG	ODN 2248/	SEQ ID NO:12
ggGGACGATCGTTGggggG	ODN 22 <i>5</i> 2	SEQ ID NO:13
ggGGAACGATCGTCgggggG	ODN <i>2</i> 253	SEQ ID NO:14
ggGGGACGATCGTCgggggG	ODM 2254	SEQ ID NO:15
ggGGGACGATCGTCGgggggG	ODN 2255	SEQ ID NO:16
ggGGTCATCGATGAgggggG	ØDN 2260	SEQ ID NO:17
ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
ggGGTCGTTCGAACGAgggggG	ODN 2294	SEQ ID NO:19
ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20
ggGGAACGACGTCGTTgggggG	ODN 2297	SEQ ID NO:21
ggGGAACGTACGTCgggggG	ODN 2298	SEQ ID NO:22
ggGGAACGTACGTACGTTgggggG /	ODN 2299	SEQ ID NO:23
ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
ggGGTCGACGTACGTCGAggggggG/	ODN 2301	SEQ ID NO:25
ggGGACCGGTACCGGTgggggG /	ODN 2302	SEQ ID NO:26
ggGTCGACGTCGAgggggG /	ODN 2303	SEQ ID NO:27
ggGGTCGACGTCGagggg /	ODN 2304	SEQ ID NO:28
ggGGAACGTTAACGTTggggģG	ODN 2305	SEQ ID NO:29
ggGGACGTCGACGTggggG'	ODN 2306	SEQ ID NO:30
ggGGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
ggTCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
ggGGACGATCGTCGgggggG	ODN 2332	SEQ ID NO:35
ggGGTCGACGTCGACgggggG	ODN 2334	SEQ ID NO:36, and
ggGGACGACGTCGTGgggggG	ODN 2336	SEQ ID NO:37,

- wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.
 - 20. The improvement of claim 1, further comprising co-administering GM-CSF to the subject.
 - 21. The improvement of claim 1, wherein the subject has a condition selected from the group consisting of a proliferative disorder and a viral infection.

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- 22. The improvement of claim 1, wherein the subject has a proliferative disorder selected from the group consisting of: hairy cell leukemia, chronic myelogenous leukemia, cutaneous T-cell leukemia, multiple myeloma, follicular lymphoma, malignant melanoma, squamous cell carcinoma, AIDS-related Kaposi's sarcoma, renal cell carcinoma, prostate carcinoma, bladder cell carcinoma, cervical dysplasia, and colon carcinoma.
- 23. The improvement of claim 1, wherein the subject has a viral infection selected from the group consisting of: hepatitis B, hepatitis C, condyloma acuminatum, human immunodeficiency virus, herpes, cytomegalovirus, Epstein-Barr virus, and papillomavirus.
- 24. A method of supplementing IFN-α treatment of a subject comprising administering to a subject in need of IFN-α treatment an effective amount of IFN-α and an isolated immunostimulatory nucleic acid.
- 25. The method of claim 24, wherein the IFN- α is administered at a dose below the clinically established effective dose for IFN- α alone.
- 26. The method of claim 24, wherein the IFN-α is administered at the maximum tolerated dose for IFN-α in the absence of the immunostimulatory nucleic acid.
 - 27. The method of claim 24, wherein the IFN-α is administered at least 20 percent below the maximum tolerated dose of IFN-α in the subject.
 - 28. The method of claim 24, wherein the IFN- α is administered at least 30 percent below the maximum tolerated dose of IFN- α in the subject.
 - 29. The method of claim 24, wherein the IFN- α is administered at least 40 percent below the maximum tolerated dose of IFN- α in the subject.

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- 30. The method of claim 24, wherein the IFN- α is administered at least 50 percent below the maximum tolerated dose of IFN- α in the subject.
- 31. The method of claim 24, wherein the immunostimulatory nucleic acid is modified.

32. The method of claim 24, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.

33. The method of claim 24, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.

- 34. The method of claim 24, wherein the immunostimulatory nucleic acid is not a palindrome.
- 35. The method of claim 24, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
- 20 36. The method of claim 24, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.
 - 37. The method of claim 36, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
 - 38. The method of claim 36, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.
- The method of claim 24, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic/acids, T-rich nucleic acids, and poly-G nucleic acids.

- 40. The method of claim 24, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.
- 41. The method of claim 24, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.
- 42. The method of claim 24, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

			/	
	ggGGTCAACGTTGAgggggG	ODN 1585	SEQ'ID NO:1	
10	tcgtcgttttgtcgttt	ODN 2022	SEQ ID NO:2	
	ggggtcgtcgttttgggggg	ODN 2184	SEQ ID NO:3	
	tcgtcgttttgtcgttttgggggg	ODN 2185 /	SEQ ID NO:4	
	ggggtcgacgtcgaggggg	ODN 2192/	SEQ ID NO:5	
	ggggtcatcgatgaggggg	ODN 2204	SEQ ID NO:6	
15	ggGGACGATCGTCgggggG	ODN 2 216	SEQ ID NO:7	
	gggggtcgtacgacgggggg	ODM 2217	SEQ ID NO:8	
	ggGGACGATATCGTCgggggG	/ ODN 2245	SEQ ID NO:9	
	ggGGACGACGTCGTCgggggG /	ØDN 2246	SEQ ID NO:10	
	ggGGACGAGCTCGTCgggggG /	/DDN 2247	SEQ ID NO:11	
20	ggGGACGTACGTCgggggG	/ ODN 2248	SEQ ID NO:12	
	ggGGACGATCGTTGggggG	ODN 2252	SEQ ID NO:13	
	ggGGAACGATCGTCgggggG /	ODN 2253	SEQ ID NO:14	
	ggGGGACGATCGTCgggggG	ODN 2254	SEQ ID NO:15	
	ggGGACGATCGTCGgggggG	ODN 2255	SEQ ID NO:16	
25	ggGGTCATCGATGAgggggG /	ODN 2260	SEQ ID NO:17	
	ggGGTCGTCGACGAgggggG /	ODN 2293	SEQ ID NO:18	
	ggGGTCGTTCGAACGAgggggg	ODN 2294	SEQ ID NO:19	
	ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20	
	ggGGAACGACGTCGTTggggggG	ODN 2297	SEQ ID NO:21	
30	ggGGAACGTACGTCgggggG	ODN 2298	SEQ ID NO:22	
	ggGGAACGTACGTACGTTgggggG	ODN 2299	SEQ ID NO:23	
	ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24	
	ggGGTCGACGTACGTCGAgggggG	ODN 2301	SEQ ID NO:25	
	ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26	
35	ggGTCGACGTCGAgggggG	ODN 2303	SEQ ID NO:27	
	ggGGTCGACG/TCGagggg	ODN 2304	SEQ ID NO:28	
	ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29	
	ggGGACGTCGACGTggggG	ODN 2306	SEQ ID NO:30	
	ggGGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31	
40	ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32	
	ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33	
	ggTCG/TCGACGAGgggggG	ODN 2330	SEQ ID NO:34	
	ggGG♠CGATCGTCGgggggG	ODN 2332	SEQ ID NO:35	
	ggGG/TCGACGTCGACGTCGAGgggggG	ODN 2334	SEQ ID NO:36, and	
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ggGGACGACGTCGTGgggggG

ODN 2336 SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

- 5 43. The method of claim 24, further comprising co-administering GM-CSF to the subject.
 - 44. The method of claim 24, wherein the subject has a condition selected from the group consisting of a proliferative disorder and a viral infection.
- 10 45. The method of claim 24, wherein the subject has a proliferative disorder selected from the group consisting of: hairy cell leukemia, chronic myelogenous leukemia, cutaneous T-cell leukemia, multiple myeloma, follicular lymphoma, malignant melanoma, squamous cell carcinoma, AIDS-related Kaposi's sarcoma, renal cell carcinoma, prostate carcinoma, bladder cell carcinoma, cervical dysplasia, and colon carcinoma.

46. The method of claim 24, wherein the subject has a viral infection selected from the group consisting of: hepatitis B, hepatitis C, condyloma acuminatum, human immunodeficiency virus, herpes, cytomegalovirus, Epstein-Barr virus, and papillomavirus.

47. A method of treating a subject to activate interferon-producing cells (IPCs) of the subject comprising

isolating IPCs from a subject in need of such treatment, culturing the IPCs in vitro,

contacting the IPCs *in vitro* with an effective amount of an isolated immunostimulatory nucleic acid, and returning the contacted IPCs to the subject.

48. The method of claim 47, further comprising contacting the IPCs *in vitro* with a growth factor.

49. The method of claim 47, further comprising contacting the IPCs in vitro with IL-3.

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- 50. The method of claim 47, further comprising contacting the IPCs in vitro with GM-CSF
- 51. The method of claim 47, wherein the IPCs are cultured in vitro in the absence of IL/3.
- 52. The method of claim 47, wherein the IPCs are cultured *in vitro* in the absence of GM-CSF.
- 53. The method of claim 47, wherein the immunostimulatory nucleic acid is modified.
- 54. The method of claim 47, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.
- 55. The method of claim 47, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.
- 56. The method of claim 47, wherein the immunostimulatory nucleic acid is not a palindrome.
- 57. The method of claim 47, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
- 25 58. The method of claim 47, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.
 - 59. The method of plaim 58, wherein the non-CpG immunostimulatory nucleic acid is a Trich nucleic acid.
 - 60. The method of claim 58, wherein the non-CpG immunostimulatory nucleic acid is a poly-G rucleic acid.

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- 61. The method of claim 47, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, T-rich nucleic acids, and poly-G nucleic acids.
- 62. The method of claim 47, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.
- 63. The method of claim 47, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.

64. The method of claim 47, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

	/ \ /	
ggGGTCAACGTTGAgggggG	ODN 1585	SEQ ID NO:1
tcgtcgttttgtcgttttgtcgtt	/ /ODN 2022	SEQ ID NO:2
ggggtcgtcgttttgggggg	/ / ODN 2184	SEQ ID NO:3
tcgtcgttttgtcgttttgggggg	ODN 2185	SEQ ID NO:4
ggggtcgacgtcgaggggg	ODN 2192	SEQ ID NO:5
ggggtcatcgatgaggggg	ODN 2204	SEQ ID NO:6
ggGGACGATCGTCgggggG	ODN 2216	SEQ ID NO:7
gggggtcgtacgacgggggg /	ODN 2217	SEQ ID NO:8
ggGGACGATATCGTCgggggG	ODN 2245	SEQ ID NO:9
ggGGGACGACGTCGTCgggggG	ODN 2246	SEQ ID NO:10
ggGGGACGAGCTCGTCgggggG	ODN 2247	SEQ ID NO:11
ggGGGACGTACGTCgggggG	ODN 2248	SEQ ID NO:12
ggGGGACGATCGTTGgggggG	ODN 2252	SEQ ID NO:13
ggGGAACGATCGTCgggggG	ODN 2253	SEQ ID NO:14
ggGGGGACGATCG/fCgggggG	ODN 2254	SEQ ID NO:15
ggGGGACGATCG/TCGgggggG	ODN 2255	SEQ ID NO:16
ggGGGTCATCGATGAgggggG	ODN 2260	SEQ ID NO:17
ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
ggGGTCGTTCGAACGAgggggG	ODN 2294	SEQ ID NO:19
ggGGACGT/TCGAACGTgggggG	ODN 2295	SEQ ID NO:20
ggGGAAÇGACGTCGTTgggggG	ODN 2297	SEQ ID NO:21
ggGGAACGTACGTCgggggG	ODN 2298	SEQ ID NO:22
ggGGAACGTACGTACGTTgggggg		SEQ ID NO:23
ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
ggGGTCGACGTACGTCGAgggggC		SEQ ID NO:25
ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
ggGTCGACGTCGAgggggG	ODN 2303	SEQ ID NO:27
ggGTCGACGTCGagggg	ODN 2304	SEQ ID NO:28
ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29

ggGGACGTCGACGTggggG		ODN 2306	SEQ ID NO:30
ggGGTCGTTCGTTgggggG		ODN 2311	SEQ ID NO:31
ggGACGATCGTCGgggggG		ODN 2328	SEQ ID NO:32
ggGTCGTCGACGAggggggG /		ODN 2329	SEQ ID NO:33
ggTCGTCGACGAGgggggG	\	ODN 2330	SEQ ID NO:34
ggGGACGATCGTCGgggggG	1	ODN 2332	SEQ ID NO:35
ggGGTCGACGTCGACGTCGAG	ggg g G	ODN 2334	SEQ ID NO:36, and
ggGGACGACGTCGTGgggggG \	$\overline{}$	ODN 2336	SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

65. A method of increasing efficacy of IFN-α treatment of a subject, comprising: administering to a subject in need of treatment with IFN-α a pharmaceutical composition comprising IFN-α, and

coadministering to the subject in need of such treatment a pharmaceutical composition comprising an immunostimulatory nucleic acid in an amount which, together with the administered IFN- α , is an effective IFN- α treatment, wherein the efficacy of the IFN- α treatment is greater than the efficacy of administering the same amount of IFN- α in the absence of coadministering the immunostimulatory nucleic acid.

66. The method of claim 65, wherein the pharmaceutical composition comprising are immunostimulatory nucleic acid is administered locally.

67. The method of claim 65, wherein the immunostimulatory nucleic acid is modified.

68. The method of claim 65, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.

69. The method of claim 65, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.

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- 70. The method of claim 65, wherein the immunostimulatory nucleic acid is not a palindrome.
- 71. The method of claim 65, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
- 72. The method of claim 65, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.
- 73. The method of claim 72, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
 - 74. The method of claim 72, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.
 - 75. The method of claim 65, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, T-rich nucleic acids, and poly-G nucleic acids.
- 76. The method of claim 65, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length
 - 77. The method of claim 65, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.

78. The method of claim 65, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

	ggGGTCAAÇGTTGAgggggG	ODN 1585	SEQ ID NO:1
	tegtegttttgtegttttgtegtt	ODN 2022	SEQ ID NO:2
30	ggggtcgtcgttttgggggg	ODN 2184	SEQ ID NO:3
	tcgtcgttttgtcgttttgggggg	ODN 2185	SEQ ID NO:4
	ggggtcgacgtcgaggggg	ODN 2192	SEQ ID NO:5
	ggggtcatcgatgaggggg	ODN 2204	SEQ ID NO:6
	ggGGGACGATCGTCgggggG	ODN 2216	SEQ ID NO:7
35	gggggtcgtacgacgggggg	ODN 2217	SEQ ID NO:8

			/
	ggGGGACGATATCGTCgggggG	ODN 2245	SEQ ID NO:9
	ggGGGACGACGTCGTCgggggG	ODN 2246	SEQ ID NO:10 /
	ggGGGACGAGCTCGTCgggggG	ODN 2247	SEQ ID NO:11/
	ggGGGACGTACGTCgggggG	ODN 2248	SEQ ID NO: J/2
5	ggGGGACGATCGTTGggggG	ODN 2252	SEQ ID NO:13
	ggGGAACGATCGTCgggggG	ODN 2253	SEQ ID NO:14
	ggGGGACGATCGTCgggggG	ODN 2254	SEQ ID,NO:15
	ggGGGACGATCGTCGgggggG	ODN 2255	SEQ ID NO:16
	ggGGTCATCGATGAgggggG	ODN 2260	SEQ/ID NO:17
10	ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
	ggGGTCGTTCGAACGAgggggG	ODN 2294	SEQ ID NO:19
	ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20
	ggGGAACGACGTCGTTgggggG	ODN 2297 /	SEQ ID NO:21
	ggGGAACGTACGTCgggggG	ODN 2298/	SEQ ID NO:22
15	ggGGAACGTACGTACGTTgggggG	ODN 2299	SEQ ID NO:23
	ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
	ggGGTCGACGTACGTCGAgggggG	ODN/2301	SEQ ID NO:25
	ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
	ggGTCGACGTCGAgggggG	ODN 2303	SEQ ID NO:27
20	ggGGTCGACGTCGagggg	ÓDN 2304	SEQ ID NO:28
	ggGGAACGTTAACGTTgggggG / `	│	SEQ ID NO:29
	ggGGACGTCGACGTggggG /	ODN 2306	SEQ ID NO:30
	ggGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
	ggGACGATCGTCGgggggG /	ODN 2328	SEQ ID NO:32
25	ggGTCGTCGACGAggggggG //	ODN 2329	SEQ ID NO:33
	ggTCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
	ggGGACGATCGTCGgggggG /	ODN 2332	SEQ ID NO:35
	ggGGTCGACGTCGACgggggG	ODN 2334	SEQ ID NO:36, and
	ggGGACGACGTCGTGgggggG /	ODN 2336	SEQ ID NO:37,
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wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

- 79. The method of claim 65, wherein the subject has a condition selected from the group consisting of a proliferative disorder and a viral infection.
- 80. The method of claim 65, wherein the subject has a proliferative disorder selected from the group consisting of: hairy cell leukemia, chronic myelogenous leukemia, cutaneous T-cell leukemia, multiple myeloma, follicular lymphoma, malignant melanoma, squamous cell carcinoma, AIDS-related Kaposi's sarcoma, renal cell carcinoma, prostate carcinoma, bladder cell carcinoma, cervical dysplasia, and colon carcinoma.

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81. The method of claim 65, wherein the subject has a viral infection selected from the group consisting of: hepatitis B, hepatitis C, condyloma acuminatum, human immunodeficiency virus, herpes, cytomegalovirus, Epstein-Barr virus, and papillomavirus.

A method of decreasing a dose of IFN- α effective for treating a subject, comprising: administering to a subject in need of treatment with IFN- α a pharmaceutical composition comprising IFN- α , and

coadministering to the subject in need of such treatment a pharmaceutical composition comprising an immunostimulatory nucleic acid in an amount which, together with the administered IFN- α , is an effective IFN- α treatment, and, wherein the amount of administered IFN- α is less than an amount of IFN- α required in the absence of coadministering the immunostimulatory nucleic acid.

- 83. The method of claim 82, wherein the amount of administered IFN-α is at least 20 percent below the amount of IFN-α required in the absence of coadministering the immunostimulatory nucleic acid.
- The method of claim 82, wherein the amount of administered IFN-α is at least 30
 percent below the amount of IFN-α required in the absence of coadministering the immunostimulatory nucleic acid.
 - 85. The method of claim 82, wherein the amount of administered IFN-α is at least 40 percent below the amount of IFN-α required in the absence of coadministering the immunostimulatory nucleic acid.
 - 86. The method of claim 82, wherein the amount of administered IFN-α is at least 50 percent below the amount of IFN-α required in the absence of coadministering the immunostimulatory nucleic acid.

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- 87. The method of claim 82, wherein the pharmaceutical composition comprising an immunostimulatory nucleic acid is administered locally.
- 88. The method of claim 82, wherein the immunostimulatory nucleic acid is modified.
- 89. The method of claim 82, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.
- 90. The method of claim 82, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.
- 91. The method of claim 82, wherein the immunostimulatory nucleic acid is not a palindrome.
- 92. The method of claim 82, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
- 20 93. The method of claim \$2, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.
 - 94. The method of claim 93, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
 - 95. The method of claim 93, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.
- 96. The/method of claim 82, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, Truch nucleic acids, and poly-G nucleic acids.

- 97. The method of claim 82, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.
- 98. The method of claim 82, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.
- 99. The method of claim 82, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

	ggGGTCAACGTTGAgggggG	ODN 1585	SÉQ ID NO:1
ı	tcgtcgttttgtcgttt	ODN 2022	∕SEQ ID NO:2
	ggggtcgtcgttttgggggg	ODN 2184 /	SEQ ID NO:3
	tcgtcgttttgtcgttttgggggg	ODN 2188	SEQ ID NO:4
	ggggtcgacgtcgagggggg	ODN 21/92	SEQ ID NO:5
	ggggtcatcgatgaggggg	ODN⁄2204	SEQ ID NO:6
	ggGGGACGATCGTCgggggG	ODN 2216	SEQ ID NO:7
	gggggtcgtacgacgggggg / \	ØDN 2217	SEQ ID NO:8
	ggGGGACGATATCGTCgggggG /	ODN 2245	SEQ ID NO:9
	ggGGGACGACGTCGTCgggggG //	ODN 2246	SEQ ID NO:10
	ggGGGACGAGCTCGTCgggggG \/	ODN 2247	SEQ ID NO:11
	ggGGGACGTACGTCgggggG	ODN 2248	SEQ ID NO:12
	ggGGGACGATCGTTGggggG /	ODN 2252	SEQ ID NO:13
	ggGGAACGATCGTCgggggG /	ODN 2253	SEQ ID NO:14
	ggGGGACGATCGTCgggggG /	ODN 2254	SEQ ID NO:15
	ggGGGACGATCGTCGgggggG /	ODN 2255	SEQ ID NO:16
	ggGGGTCATCGATGAgggggG/	ODN 2260	SEQ ID NO:17
	ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
	ggGGTCGTTCGAACGAgggggG	ODN 2294	SEQ ID NO:19
	ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20
	ggGGAACGACGTCGTTggggggG	ODN 2297	SEQ ID NO:21
ı	ggGGAACGTACGTØgggggG	ODN 2298	SEQ ID NO:22
	ggGGAACGTACGTACGTTgggggG	ODN 2299	SEQ ID NO:23
	ggGGTCACCGG/TGAgggggG	ODN 2300	SEQ ID NO:24
	ggGGTCGACG/TACGTCGAgggggG	ODN 2301	SEQ ID NO:25
	ggGGACCGG/TACCGGTgggggG	ODN 2302	SEQ ID NO:26
	ggGTCGACGTCGAgggggG	ODN 2303	SEQ ID NO:27
	ggGGTCGACGTCGagggg	ODN 2304	SEQ ID NO:28
	ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29
	ggGGACGTCGACGTggggG	ODN 2306	SEQ ID NO:30
	ggGGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
ı	ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
	ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
	gg/TCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
	ggGGACGATCGTCGgggggG	ODN 2332	SEQ ID NO:35
	\(\frac{1}{2} \text{gGGTCGACGTCGACggggggG} \)	ODN 2334	SEQ ID NO:36, and

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ggGGACGACGTCGTGgggggG

ODN 2336 SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

- 5 100. The method of claim 82, wherein the subject has a condition selected from the group consisting of a proliferative disorder and a viral infection.
 - 101. The method of claim 82, wherein the subject has a proliferative disorder selected from the group consisting of: hairy cell leukemia, chronic myelogenous leukemia, cutaneous T-cell leukemia, multiple myeloma, follicular lymphoma, malignant melanoma, squamous cell carcinoma, AIDS-related Kaposi's sarcoma, renal cell carcinoma, prostate carcinoma, bladder cell carcinoma, cervical dysplasia, and colon carcinoma.
 - 102. The method of claim 82, wherein the subject has a viral infection selected from the group consisting of: hepatitis B, hepatitis C, condyloma acuminatum, human immunodeficiency virus, herpes, cytomegalovirus, Epstein-Barr virus, and papillomavirus.
 - 103. A method of preventing an IFN-α treatment-related side effect in a subject receiving or in need of treatment with IFN-α, comprising

administering to a subject in need of treatment with IFN- α a pharmaceutical composition comprising IFN- α , and

coadministering to the subject in need of such treatment a pharmaceutical composition comprising an immunostimulatory nucleic acid in an amount which, together with the administered IFN- α is an effective IFN- α treatment, and, wherein an IFN- α treatment-related side effect is reduced in comparison to the side effect when IFN- α is administered in the absence of coadministering the immunostimulatory nucleic acid.

- 104. The method of claim 103, wherein the pharmaceutical composition comprising an immunostimulatory nucleic acid is administered locally.
- 105. The method of claim 103, wherein the IFN- α treatment-related side effect is systemic.

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- 106. The method of claim 103, wherein the IFN-α treatment-related side effect is selected from the group consisting of flu-like syndrome, fever, headache, chills, myalgia, fatigue, anorexia, nausea, vomiting, diarrhea, and depression.
- 107. The method of claim 103, wherein the immunostimulatory nucleic acid is modified.
- 108. The method of claim 103, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.
- 109. The method of claim 103, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.
- 110. The method of claim 103, wherein the immunostimulatory nucleic acid is not a palindrome.
- 111. The method of claim 103, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
- 112. The method of claim 103, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.
- 25 113. The method of claim 112, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
 - 114. The method of claim 112, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.

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- 115. The method of claim 103, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, T-rich nucleic acids, and poly-G nucleic acids.
- 5 116. The method of claim 103, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.
 - 117. The method of claim 103, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.

118. The method of claim 103, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

	/	
ggGGTCAACGTTGAgggggG	ODN 1585	SEQ ID NO:1
tcgtcgttttgtcgttttgtcgtt	ODN 2022	SEQ ID NO:2
ggggtcgtcgttttgggggg	ØDN 2184	SEQ ID NO:3
tcgtcgttttgtcgttttgggggg	/ \/ODN 2185	SEQ ID NO:4
ggggtcgacgtcgaggggg	/ / ODN 2192	SEQ ID NO:5
ggggtcatcgatgaggggg	// ODN 2204	SEQ ID NO:6
ggGGGACGATCGTCgggggG	ODN 2216	SEQ ID NO:7
gggggtcgtacgacggggg	ODN 22/17	SEQ ID NO:8
ggGGACGATATCGTCgggggG	ODN 2245	SEQ ID NO:9
ggGGACGACGTCGTCgggggG /	ODN 2246	SEQ ID NO:10
ggGGACGAGCTCGTCgggggG /	ODN 2247	SEQ ID NO:11
ggGGGACGTACGTCgggggG /	ODN 2248	SEQ ID NO:12
ggGGGACGATCGTTGggggG /	ODN 2252	SEQ ID NO:13
ggGGAACGATCGTCgggggG /	ODN 2253	SEQ ID NO:14
ggGGGACGATCGTCgggggG	ODN 2254	SEQ ID NO:15
ggGGGACGATCGTCGgggggG	ODN 2255	SEQ ID NO:16
ggGGGTCATCGATGAgggggG	ODN 2260	SEQ ID NO:17
ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
ggGGTCGTTCGAAC&AgggggG	ODN 2294	SEQ·ID NO:19
ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20
ggGGAACGACGT/ĆGTTgggggG	ODN 2297	SEQ ID NO:21
ggGGAACGTAÇGTCgggggG	ODN 2298	SEQ ID NO:22
ggGGAACGTACGTTgggggG	ODN 2299	SEQ ID NO:23
ggGGTCACCØGTGAgggggG	ODN 2300	SEQ ID NO:24
ggGGTCGA¢GTACGTCGAgggggG	ODN 2301	SEQ ID NO:25
ggGGACCØGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
ggGTCGA/CGTCGAgggggG	ODN 2303	SEQ ID NO:27
ggGGTCGACGTCGagggg	ODN 2304	SEQ ID NO:28
ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29
ggGGACGTCGACGTggggG	ODN 2306	SEQ ID NO:30
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ggGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
ggTCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
ggGGACGATCGTCGgggggG	ODN 2332	SEQ ID NO:35
ggGGTCGACGTCGACgggggG	ODN 2334	SEQ ID NO:36, and
ggGGACGACGTCGTGgggggG	ODN 2336	SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

- 119. The method of claim 103, wherein the subject has a condition selected from the group consisting of a proliferative disorder and a viral infection.
- 120. The method of claim 103, wherein the subject has a proliferative disorder selected from the group consisting of: hairy cell leukemia, chronic myelogenous leukemia, cutaneous T-cell leukemia, multiple myeloma, follicular lymphoma, malignant melanoma, squamous cell carcinoma, AIDS-related Kaposi's sarcoma, renal cell carcinoma, prostate carcinoma, bladder cell carcinoma, cervical dysplasia, and colon carcinoma.

121. The method of claim 103, wherein the subject has a viral infection selected from the group consisting of: hepatitis B, hepatitis C, condyloma acuminatum, human immurodeficiency virus, herpes, cytomegalovirus, Epstein-Barr virus, and papillomavirus.

122. A method of enhancing efficacy of IFN-α treatment in a subject in need of such treatment, comprising

administering to a subject in need of such treatment an amount of a pharmaceutical composition comprising IFN- α effective for treating a condition of the subject;

isolating natural interferon-producing cells (IPCs) from a donor;

contacting the isolated IPCs $ex\ vivo$ with an amount of a pharmaceutical composition comprising an immunostimulatory nucleic acid effective for inducing the IPCs to release IFN- α ; and

administering the contacted cells to the subject.

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- 123. The method of claim 122, wherein the donor is the subject.
- 124. The method of claim 122 further comprising contacting the isolated IPCs with an antigen.
 - 125. The method of claim 122, wherein the administering the contacted cells comprises local injection.
- 126. The method of claim 125, wherein the local injection is via a blood vessel supplying a target tissue.
 - 127. The method of claim 126, wherein the blood vessel is selected from the group consisting of a hepatic artery, a portal vein, a celiac artery, and a splenic artery.
 - 128. The method of claim 122, wherein the immunostimulatory nucleic acid is modified.
 - 129. The method of claim 122, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.
 - 130. The method of claim 122, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.
 - 131. The method of claim 122, wherein the immunostimulatory nucleic acid is not a palindrome.
 - 132. The method of claim 122, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.

- 133. The method of claim 122, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.
- 134. The method of claim 133, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
 - 135. The method of claim 133, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.
- 136. The method of claim 122, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, T-rich nucleic acids, and poly-G nucleic acids.
 - 137. The method of claim 122, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.
 - 138. The method of claim 122, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.
- 20 139. The method of claim 122, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

	ggGGTCAACGTTGAgggggG	ODN 1585	SEQ ID NO:1
	tcgtcgttttgtcgttttgtcgtt /	ODN 2022	SEQ ID NO:2
	ggggtcgtcgttttgggggg /	ODN 2184	SEQ ID NO:3
25	tcgtcgttttgtcgttttgggggg /	ODN 2185	SEQ ID NO:4
	ggggtcgacgtcgagggggg/	ODN 2192	SEQ ID NO:5
	ggggtcatcgatgaggggg	ODN 2204	SEQ ID NO:6
	ggGGGACGATCG/fCgggggG	ODN 2216	SEQ ID NO:7
	gggggtcgtacgacggggg	ODN 2217	SEQ ID NO:8
30	ggGGGACGATÁTCGTCgggggG	ODN 2245	SEQ ID NO:9
	ggGGGACGACGTCGTCgggggG	ODN 2246	SEQ ID NO:10
	ggGGGACGAGCTCGTCgggggG	ODN 2247	SEQ ID NO:11
	ggGGGACGTACGTCgggggG	ODN 2248	SEQ ID NO:12
	ggGGGAGGATCGTTGggggG	ODN 2252	SEQ ID NO:13
35	ggGGAACGATCGTCgggggG	ODN 2253	SEQ ID NO:14
	ggGGGGACGATCGTCgggggG	ODN 2254	SEQ ID NO:15
	ggGGGACGATCGTCGgggggG	ODN 2255	SEQ ID NO:16
	ggGGGTCATCGATGAgggggG	ODN 2260	SEQ ID NO:17
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ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
ggGGTCGTTCGAACGAgggggG	ODN 2294	SEQ ID NO:19
ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20/
ggGGAACGACGTCGTTgggggG	ODN 2297	SEQ ID NO:21
ggGGAACGTACGTCgggggG	ODN 2298	SEQ ID NO;22
ggGGAACGTACGTACGTTgggggG	ODN 2299	SEQ ID NO:23
ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
ggGGTCGACGTACGTCGAgggggG	ODN 2301	SEQ ID/NO:25
ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ĮĎ NO:26
ggGTCGACGTCGAgggggG	ODN 2303	SEQ'ÍD NO:27
ggGGTCGACGTCGagggg	ODN 2304	SĘQ ID NO:28
ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29
ggGGACGTCGACGTggggG	ODN 2306 /	SEQ ID NO:30
ggGGTCGTTCGTTgggggG	ODN 2311/	SEQ ID NO:31
ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
ggGTCGTCGACGAggggggG	ODN 23⁄29	SEQ ID NO:33
ggTCGTCGACGAGgggggG	ODN 2 330	SEQ ID NO:34
ggGGACGATCGTCGgggggG	ODM 2332	SEQ ID NO:35
ggGGTCGACGTCGACggggggG	OØN 2334	SEQ ID NO:36, and
ggGGACGACGTCGTGgggggG	ØDN 2336	SEQ ID NO:37,
/	/	

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

- 140. The method of claim 122, wherein the subject has a condition selected from the group consisting of a proliferative disorder and a viral infection.
- 141. The method of claim 122, wherein the subject has a proliferative disorder selected from the group consisting of: hairy cell leukemia, chronic myelogenous leukemia, cutaneous T-cell leukemia, multiple myeloma, follicular lymphoma, malignant melanoma, squamous cell carcinoma, AIDS-related Kaposi's sarcoma, renal cell carcinoma, prostate carcinoma, bladder cell carcinoma, cervical dysplasia, and colon carcinoma.
- 142. The method of claim 122, wherein the subject has a viral infection selected from the group consisting of: hepatitis B, hepatitis C, condyloma acuminatum, human immunodeficiency virus, herpes, cytomegalovirus, Epstein-Barr virus, and papillomavirus.

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143. A method of supporting survival of natural interferon-producing cells (IPCs) *in vitro*, comprising

isolating IPCs from a subject;

culturing the IPCs in a sterile medium suitable for tissue culture; and contacting the IPCs *in vitro* with an amount of immunostimulatory nucleic acid effective to support the growth of the IPCs in the absence of interleukin 3 (IL-3).

- 144. The method of claim 143, wherein the IPCs are precursor type 2 dendritic cells (pDC2s).
- 145. The method of claim 143, wherein the IPCs are cultured in the absence of IL-3.
- 146. The method of claim 143, wherein the IPCs are cultured in the absence of GM-CSF.
- 15 147. The method of claim 143, wherein the immunostimulatory nucleic acid is modified.
 - 148. The method of claim 143, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.
 - 149. The method of claim 143, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.
- 25 150. The method of claim 143, wherein the immunostimulatory nucleic acid is not a palindrome.
 - 151. The method of claim 143, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
 - 152. The method of claim 143, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.

- 153. The method of claim 152, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
- 5 154. The method of claim 152, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.
 - 155. The method of claim 143, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, T-rich nucleic acids, and poly-G nucleic acids.
 - 156. The method of claim 143, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.
- 15 157. The method of claim 143, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.
 - 158. The method of claim 143, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

20	ggGGTCAACGTTGAgggggG/	ODN 1585	SEQ ID NO:1
	tegtegttttgtegttttgtegtt	ODN 2022	SEQ ID NO:2
	ggggtcgtcgttttgggggg	ODN 2184	SEQ ID NO:3
	tcgtcgttttgtcgttttgggggg	ODN 2185	SEQ ID NO:4
	ggggtcgacgtcgagggggg/	ODN 2192	SEQ ID NO:5
25	ggggtcatcgatgagggggg	ODN 2204	SEQ ID NO:6
	ggGGGACGATCG/TCgggggG	ODN 2216	SEQ ID NO:7
	gggggtcgtacgacgggggg	ODN 2217	SEQ ID NO:8
	ggGGACGAT/ATCGTCgggggG	ODN 2245	SEQ ID NO:9
	ggGGGACGACGTCGTCgggggG	ODN 2246	SEQ ID NO:10
30	ggGGGACQAGCTCGTCgggggG	ODN 2247	SEQ ID NO:11
	ggGGGAÇGTACGTCgggggG	ODN 2248	SEQ ID NO:12
	ggGGGACGATCGTTGggggG	ODN 2252	SEQ ID NO:13
	ggGGAACGATCGTCgggggG	ODN 2253	SEQ ID NO:14
	ggGGGGACGATCGTCgggggG	ODN 2254	SEQ ID NO:15
35	ggGGGACGATCGTCGgggggG	ODN 2255	SEQ ID NO:16
	ggGGTCATCGATGAgggggG	ODN 2260	SEQ ID NO:17
	ggGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
	ggGGTCGTTCGAACGAgggggG	ODN 2294	SEQ ID NO:19
	ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20
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	ggGGAACGACGTCGTTgggggG	ODN 2297	SEQ ID NO:21
	ggGGAACGTACGTCgggggG	ODN 2298	SEQ ID NO:22
	ggGGAACGTACGTACGTTgggggG	ODN 2299	SEQ ID NO:23
	ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
5	ggGGTCGACGTACGTCGAgggggG	ODN 2301	SEQ ID NO:25
	ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
	ggGTCGACGTCGAgggggG	ODN 2303	SEQ ID NO:27
	ggGGTCGACGTCGagggg	ODN 2304	SEQ ID NO:28
	ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29
)	ggGGACGTCGACGTggggG	ODN 2306	SEQ ID NO:30
	ggGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
	ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
	ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
	ggTCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
5	ggGGACGATCGTCGgggggG	ODN 2332	SEQ ID NO:35
	ggGGTCGACGTCGACGTCGAGgggggG	ODN 2334	SEQ ID NO:36, and
	ggGGACGACGTCGTGgggggG	ODN 2336	SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

159. A method of stimulating isolated interferon-producing cells (IPCs) *in vitro*, comprising

isolating IPCs from a subject;

culturing the IPCs in a sterile medium suitable for tissue culture; and contacting the IPCs *in vitro* with an amount of immunostimulatory nucleic acid effective to induce secretion of at least one type I interferon.

- 160. The method of claim 159, wherein the IPCs are precursor type 2 dendritic cells (pDC2s).
- 161. The method of claim 159, wherein the type I interferon is an IFN- α .
- 162. The method of claim 159, wherein the IPCs are cultured in the absence of IL-3.
- The method of claim 159, wherein the IPCs are cultured in the absence of GM-CSF.
 - 164. The method of claim 159, wherein the immunostimulatory nucleic acid is modified.

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- 165. The method of claim 159, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.
- 166. The method of claim 159, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.
- 167. The method of claim 159, wherein the immunostimulatory nucleic acid is not a palindrome.
 - 168. The method of claim 159, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
- 15 169. The method of claim 159, wherein the immunostimulatory nucleic acid is a non-CpG nucleic acid.
 - 170. The method of claim 169, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
 - 171. The method of claim 169, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.
 - 172. The method of claim 159, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, 7-rich nucleic acids, and poly-G nucleic acids.
 - 173. The method of claim 159, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.
 - 174. The method of claim 159, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.

175. The method of claim 159, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

	ggGGTCAACGTTGAgggggG	ODN 1585	SEQ ID NO:1
5	tcgtcgttttgtcgttt	ODN 2022	SEQ ID NO:2
	ggggtcgtcgttttgggggg	ODN 2184	SEQ JØ NO:3
	tcgtcgttttgtcgttttgggggg	ODN 2185	SEQ ID NO:4
	ggggtcgacgtcgaggggg	ODN 2192	SEQ ID NO:5
	ggggtcatcgatgaggggg	ODN 2204 /	SEQ ID NO:6
10	ggGGACGATCGTCgggggG	ODN 2216/	SEQ ID NO:7
	gggggtcgtacgacgggggg	ODN 221/7	SEQ ID NO:8
	ggGGACGATATCGTCgggggG	ODN 2 245	SEQ ID NO:9
	ggGGACGACGTCGTCgggggG	ODN 2246	SEQ ID NO:10
	ggGGACGAGCTCGTCgggggG	ODN 2247	SEQ ID NO:11
1.5	ggGGGACGTACGTCgggggG	ÓDN 2248	SEQ ID NO:12
	ggGGACGATCGTTGggggG	/ ODN 2252	SEQ ID NO:13
	ggGGAACGATCGTCgggggG / \	ODN 2253	SEQ ID NO:14
	ggGGGACGATCGTCgggggG /	ODN 2254	SEQ ID NO:15
	ggGGGACGATCGTCGgggggG //	ODN 2255	SEQ ID NO:16
20	ggGGTCATCGATGAgggggG ///	ODN 2260	SEQ ID NO:17
	ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
	ggGGTCGTTCGAACGAgggggG/	ODN 2294	SEQ ID NO:19
	ggGGACGTTCGAACGTgggggg	ODN 2295	SEQ ID NO:20
	ggGGAACGACGTCGTTggggggG	ODN 2297	SEQ ID NO:21
25	ggGGAACGTACGTCggggggG	ODN 2298	SEQ ID NO:22
	ggGGAACGTACGTACGTTgggggG	ODN 2299	SEQ ID NO:23
	ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
	ggGGTCGACGTACG/TCGAgggggG	ODN 2301	SEQ ID NO:25
	ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
30	ggGTCGACGTCG⁄AgggggG	ODN 2303	SEQ ID NO:27
	ggGGTCGACGT/CGagggg	ODN 2304	SEQ ID NO:28
	ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29
	ggGGACGT¢GACGTggggG	ODN 2306	SEQ ID NO:30
	ggGGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
35	ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
	ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
	ggTCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
	ggGGACGATCGTCGgggggG	ODN 2332	SEQ ID NO:35
	gg/GTCGACGTCGACGTCGAGgggggG	ODN 2334	SEQ ID NO:36, and
40	g g GGACGACGTCGTGgggggG	ODN 2336	SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

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- 176. A method of stimulating production of a plurality of type I IFN subtypes, comprising contacting IPCs with an amount of immunostimulatory nucleic acid effective to induce secretion of at least two type I interferons.
- 5 177. The method of claim 176, wherein the contacting occurs in vivo
 - 178. The method of claim 176, wherein the contacting occurs in vitro.
- 179. The method of claim 176, wherein the IPCs are precursor type 2 dendritic cells (pDC2s).
 - 180. The method of claim 176, wherein the IPO's are isolated.
 - 181. The method of claim 176, wherein the IPCs are induced to secrete at least three type I interferons.
 - 182. The method of claim 176, wherein the IPCs are induced to secrete at least four type I interferons.
- 20 183. The method of claim 176, wherein the IPCs are induced to secrete at least five type I interferons.
 - 184. The method of claim 176, wherein the IPCs are induced to secrete at least six type I interferons.
 - 185. The method of claim 176, wherein the IPCs are induced to secrete at least seven type I interferons.
 - The method of claim 176, wherein the IPCs are induced to secrete at least eight type I interferons.
 - 187. The method of claim 176, wherein the immunostimulatory nucleic acid is modified.

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- 188. The method of claim 176, wherein the immunostimulatory nucleic acid comprises a backbone with at least one nuclease-resistant internucleotide linkage selected from the group consisting of: phosphorothioate, phosphorodithioate, methylphosphonate, and peptide.
- 189. The method of claim 176, wherein the immunostimulatory nucleic acid comprises at least one nucleotide analog or derivative.
- 190. The method of claim 176, wherein the immunostimulatory nucleic acid is not a palindrome.
 - 191. The method of claim 176, wherein the immunostimulatory nucleic acid is a CpG nucleic acid.
 - 192. The method of claim 176, wherein the imphunostimulatory nucleic acid is a non-CpG nucleic acid.
 - 193. The method of claim 192, wherein the non-CpG immunostimulatory nucleic acid is a T-rich nucleic acid.
 - 194. The method of claim 192, wherein the non-CpG immunostimulatory nucleic acid is a poly-G nucleic acid.
- 25 195. The method of claim 176, wherein the immunostimulatory nucleic acid is any combination of at least two nucleic acids selected from the group consisting of: CpG nucleic acids, T-rich nucleic acids, and poly-G nucleic acids.
 - 196. The method of claim 176, wherein the immunostimulatory nucleic acid is between 8 and 100 nucleotides in length.

197. The method of claim 176, wherein the immunostimulatory nucleic acid is between 12 and 40 nucleotides in length.

198. The method of claim 176, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

ggGGTCAACGTTGAgggggG	ODN 1585	SEQ ID NO;1
tcgtcgttttgtcgttt	ODN 2022	SEQ ID NÓ:2
ggggtcgtcgttttgggggg	ODN 2184	SEQ ID/NO:3
tcgtcgttttgtcgttttgggggg	ODN 2185	SEQJID NO:4
ggggtcgacgtcgaggggg	ODN 2192	SEØ ID NO:5
ggggtcatcgatgaggggg	ODN 2204	SÉQ ID NO:6
ggGGACGATCGTCgggggG	ODN 2216 /	SEQ ID NO:7
gggggtcgtacgacggggg	ODN 2217/	SEQ ID NO:8
ggGGACGATATCGTCgggggG	ODN 2245	SEQ ID NO:9
ggGGACGACGTCGTCgggggG	ODN/2246	SEQ ID NO:10
ggGGACGAGCTCGTCgggggG	ODN 2247	SEQ ID NO:11
ggGGACGTACGTCgggggG	ØDN 2248	SEQ ID NO:12
ggGGACGATCGTTGggggG	ODN 2252	SEQ ID NO:13
ggGGAACGATCGTCgggggG / //	ODN 2253	SEQ ID NO:14
ggGGGACGATCGTCgggggG /	ODN 2254	SEQ ID NO:15
ggGGACGATCGTCGgggggG //	ODN 2255	SEQ ID NO:16
ggGGTCATCGATGAgggggG //	ODN 2260	SEQ ID NO:17
ggGGTCGTCGACGAgggggG	ODN/2293	SEQ ID NO:18
ggGGTCGTTCGAACGAgggggG /	ODN 2294	SEQ ID NO:19
ggGGACGTTCGAACGTgggggG /	ODN 2295	SEQ ID NO:20
ggGGAACGACGTCGTTggggggG	ODN 2297	SEQ ID NO:21
ggGGAACGTACGTCgggggG/	ODN 2298	SEQ ID NO:22
ggGGAACGTACGTT/gggggG	ODN 2299	SEQ ID NO:23
ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
ggGGTCGACGTACGTQGAgggggG	ODN 2301	SEQ ID NO:25
ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
ggGTCGACGTCGAgggggG	ODN 2303	SEQ ID NO:27
ggGGTCGACGTCGagggg	ODN 2304	SEQ ID NO:28
ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29
ggGGACGTCGACGTggggG	ODN 2306	SEQ ID NO:30
ggGGGTCGT/CGTTgggggG	ODN 2311	SEQ ID NO:31
ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
ggTCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
ggGGACGATCGTCGgggggG	ODN 2332	SEQ ID NO:35
ggGGT¢GACGTCGACGTCGAGgggggG	ODN 2334	SEQ ID NO:36, and
ggGGACGACGTCGTGgggggG	ODN 2336	SEQ ID NO:37,
1		,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

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- 199. A method of inhibiting IL-12 production, comprising contacting IL-12-producing cells, in the presence of interferon-producing cells under conditions in which the IL-12-producing cells normally produce IL-12, with an
- 5 immunostimulatory nucleic acid in an amount effective for inducing secretion of type I interferon.
 - 200. The method of claim 199, wherein the immunostimulatory nucleic acid has a sequence selected from the group consisting of

)	ggGGTCAACGTTGAgggggG	ODN 1585	SEQ ID NO:1
	tcgtcgttttgtcgttt	ODN 2022	SEQ ID NO:2
	ggggtcgtcgttttgggggg	ODN 2184	SEQ ID NO:3
	tcgtcgttttgtcgttttgggggg	ODN 2185	SEQ'ÍD NO:4
	ggggtcgacgtcgagggggg	ODN 2192	SEQ ID NO:5
;	ggggtcatcgatgaggggg	ODN 2204	SEQ ID NO:6
	ggGGACGATCGTCgggggG	ODN 2216 /	SEQ ID NO:7
	gggggtcgtacgacgggggg	ODN 2217	SEQ ID NO:8
	ggGGACGATATCGTCgggggG	ODN 2 2 45	SEQ ID NO:9
	ggGGACGACGTCGTCgggggG	ODN/2246	SEQ ID NO:10
)	ggGGACGAGCTCGTCgggggG	ODN 2247	SEQ ID NO:11
	ggGGGACGTACGTCgggggG	ØDN 2248	SEQ ID NO:12
	ggGGACGATCGTTGggggG	ODN 2252	SEQ ID NO:13
	ggGGAACGATCGTCgggggG / \	ODN 2253	SEQ ID NO:14
	ggGGGACGATCGTCgggggG / /	ODN 2254	SEQ ID NO:15
,	ggGGACGATCGTCGgggggG	ODN 2255	SEQ ID NO:16
	ggGGTCATCGATGAgggggG //	ODN 2360	SEQ ID NO:17
	ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
	ggGGTCGTTCGAACGAgggggG/	ODN 2294	SEQ ID NO:19
	ggGGACGTTCGAACGTggggg¢Ġ	ODN 2295	SEQ ID NO:20
)	ggGGAACGACGTCGTTgggggG	ODN 2297	SEQ ID NO:21
	ggGGAACGTACGTCgggggG	ODN 2298	SEQ ID NO:22
	ggGGAACGTACGTACGTTgggggG	ODN 2299	SEQ ID NO:23
	ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
	ggGGTCGACGTACGTCGAgggggG	ODN 2301	SEQ ID NO:25
5	ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
	ggGTCGACGTØGAgggggG	ODN 2303	SEQ ID NO:27
	ggGGTCGACGTCGagggg	ODN 2304	SEQ ID NO:28
	ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29
	ggGGAC&TCGACGTggggG	ODN 2306	SEQ ID NO:30
)	ggGGG/TCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
	ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
	ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
	ggTCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
	/ggGGACGATCGTCGggggggG	ODN 2332	SEQ ID NO:35

ggGGTCGACGTCGACGTCGAGgggggG ggGGACGACGTCGTGgggggG

ODN 2334 -ODN 2336 SEQ ID NO:36, and SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

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201. An isolated nucleic acid having a sequence selected from the group consisting of:

tcgtcgttttgtcgtt	ODN 2022	SEQ ID NO:2
ggggtcgtcgttttgggggg	ODN 2184	SEQ ID NO:3
tcgtcgttttgtcgttttgggggg	ODN 2185	SEQ ID NO:4
ggggtcgacgtcgaggggg	ODN 2192	SEQ ID NO:5
ggggtcatcgatgaggggg	ODN 2204	SEQ ID NO:6
ggGGACGATCGTCgggggG	ODN 2216	SEQ ID NO:7
gggggtcgtacgacgggggg	ODN 2217	SEQ ID NO:8
ggGGACGATATCGTCgggggG	ODN 2245	SEQ ID NO:9
ggGGGACGACGTCGTCgggggG	ODN 2246	SEQ ID NO:10
ggGGACGAGCTCGTCgggggG	ODN 2247	SEQ ID NO:11
ggGGGACGTACGTCgggggG	ODN 2248	SEQ ID NO:12
ggGGACGATCGTTGggggG	ODN 2252	SEQ ID NO:13
ggGGAACGATCGTCgggggG	ODN 2253	SEQ ID NO:14
ggGGGACGATCGTCgggggG	ODN 2254	SEQ ID NO:15
ggGGACGATCGTCGgggggG	ODN 2255	SEQ ID NO:16
ggGGGTCATCGATGAgggggG	ODN 2260	SEQ ID NO:17
ggGGTCGTCGACGAgggggG	ODN 2293	SEQ ID NO:18
ggGGTCGTTCGAACGAgggggG	ODN 2294	SEQ ID NO:19
ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20
ggGGAACGACGTCGTTgggggG	ODN 2297	SEQ ID NO:21
ggGGAACGTACGTCgggggG	ODN 2298	SEQ ID NO:22
ggGGAACGTACGTACGTTgggggG	ODN 2299	SEQ ID NO:23
ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
ggGGTCGACGTACGTCGAgggggG	ODN 2301	SEQ ID NO:25
ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
ggGTCGACGTCGAgggggG	ODN 2303	SEQ ID NO:27
ggGGTCGACGTCGagggg	ODN 2304	SEQ ID NO:28
ggGGAACGTTAACGTTgggggG	ODN 2305	SEQ ID NO:29
ggGGACGTCGACGTggggG	ODN 2306	SEQ ID NO:30
ggGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
ggTCGTCGACGAGgggggG	ODN 2330	SEQ ID NO:34
ggGGACGATCGTCGgggggG	ODN 2332	SEQ ID NO:35
ggGGTCGACGTCGACgggggG	ODN 2334	SEQ ID NO:36, and
ggGGACGACGTCGTGgggggG	ODN 2336	SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage.

202. A pharmaceutical composition comprising
an isolated nucleic acid having a sequence selected from the group consisting of:
tcgtcgttttgtcgttttgtcgtt

ODN 2022 SEQ ID NO:2

	/		
	tcgtcgttttgtcgttttgtcgtt /	ODN 2022	SEQ ID NO:2
5	ggggtcgtcgttttgggggg /	ODN 2184	SEQ ID NO:3
	tcgtcgttttgtcgttttgggggg /	ODN 2185	SEQ ID NO:4
	ggggtcgacgtcgaggggg /	ODN 2192	SEQ ID NO:5
	ggggtcatcgatgagggggg /	ODN 2204	SEQ ID NO:6
	ggGGGACGATCGTCgggggG /	ODN 2216	SEQ ID NO:7
10	gggggtcgtacgacgggggg /	ODN 2217	SEQ ID NO:8
	ggGGGACGATATCGTCgggggG /	ODN 2245	SEQ ID NO:9
	ggGGGACGACGTCGTCgggggG /	ODN 2246	SEQ ID NO:10
	ggGGACGAGCTCGTCgggggG /	ODN 2247	SEQ ID NO:11
	ggGGACGTACGTCgggggG /	ODN 2248	SEQ ID NO:12
15	ggGGACGATCGTTGggggG	ODN 2252	SEQ ID NO:13
	ggGGAACGATCGTCgggggG /	ODN 2253	SEQ ID NO:14
	ggGGGACGATCGTCgggggG /	ODN 2254	SEQ ID NO:15
	ggGGGACGATCGTCGgggggG /	ODN 2255	SEQ ID NO:16
	ggGGTCATCGATGAgggggG /	ODN 2260	SEQ ID NO:17
20	ggGGTCGTCGACGAgggggG /	ODN 2293	SEQ ID NO:18
	ggGGTCGTTCGAACGAgggggĠ	ODN 2294	SEQ ID NO:19
	ggGGACGTTCGAACGTgggggG	ODN 2295	SEQ ID NO:20
	ggGGAACGACGTCGTTgggggG	ODN 2297	SEQ ID NO:21
	ggGGAACGTACGTCgggggĢ	ODN 2298	SEQ ID NO:22
25	ggGGAACGTACGTACGTT/gggggG	ODN 2299	SEQ ID NO:23
	ggGGTCACCGGTGAgggggG	ODN 2300	SEQ ID NO:24
	ggGGTCGACGTACGTCG/AgggggG	ODN 2301	SEQ ID NO:25
	ggGGACCGGTACCGGTgggggG	ODN 2302	SEQ ID NO:26
	ggGTCGACGTCGAgggggG	ODN 2303	SEQ ID NO:27
30	ggGGTCGACGTCGaggg ^f g	ODN 2304	SEQ ID NO:28
	ggGGAACGTTAACGT/TggggggG	ODN 2305	SEQ ID NO:29
	ggGGACGTCGACGTggggG	ODN 2306	SEQ ID NO:30
	ggGGTCGTTCGTTgggggG	ODN 2311	SEQ ID NO:31
	ggGACGATCGTCGgggggG	ODN 2328	SEQ ID NO:32
35	ggGTCGTCGACGAggggggG	ODN 2329	SEQ ID NO:33
	ggTCGTCGACGAG/ggggggG	ODN 2330	SEQ ID NO:34
	ggGGACGATCGT¢GgggggG	ODN 2332	SEQ ID NO:35
	ggGGTCGACGTCGAGgggggG	ODN 2334	SEQ ID NO:36, and
	ggGGACGACGT ¢ GTGgggggG	ODN 2336	SEQ ID NO:37,

wherein each lower case letter represents phosphorothioate linkage and each upper case letter indicates phosphodiester linkage; and

a pharmaceutically acceptable carrier.

203. The pharmaceutical composition of claim 202, further comprising IFN-α.